Amendment Under 37 C.F.R. § 1.111 Attorney docket: Q63915 Application No. 09/820,843

AMENDMENTS TO THE CLAIMS

This listing of claims will replace all prior versions and listings of claims in the

application:

LISTING OF CLAIMS:

1.-19. (Canceled).

20. (Currently amended) A method for identifying a candidate protein useful as an

anti-infective, comprising:

(a) calculating computationally protein sequence-based attributes from protein sequences

of a pathogenic organism, wherein said protein sequences are predicted either from whole

genomic sequences, or are predicted-from partial genomic sequences comprising at least one

chromosome, and wherein said protein sequence-based attributes comprise: percentage of

charged amino acids, percentage hydrophobicity, distance of protein sequence from a fixed

reference frame, measure of dipeptide complexity, and measure of hydrophobicity from a fixed

reference frame;

(b) clustering computationally said protein sequences based on said protein sequence-

based attributes using Principle Component Analysis, and displaying the results of said

clustering;

(c) identifying computationally outlier proteins, wherein said outlier proteins appear

outside a main cluster;

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- (d) comparing <u>said</u> outlier proteins with protein sequences in databases to identify outlier <u>proteins</u> that are unique to said pathogenic organism, and to identify outlier proteins that are <u>homologous</u> or identical to proteins known to be involved in virulence;
- (e) selecting an outlier protein identified in step (d) for further testing as an anti-infective; and
 - (f) validating the said outlier protein selected in step (e) as an anti-infective.
- 21. (Previously presented) The method of claim 20, wherein said pathogenic organism is selected from the group consisting of B.burgdorfei, C.jejuni, C.pneumoniae, C.trachomatis, H.influenzae, H.pylori, L.major, M.genitalium, M.pneumoniae, M.tuberculosis, N.menigitis, P.aeruginosa, P.falciparum, R.prowazekii, T.pallidum, and V.cholerae.
- 22. (Previously presented) The method of claim 20, wherein said protein sequencebased attributes comprise fixed protein attributes and variable protein attributes.
- 23. (Previously presented) The method of claim 22, wherein a variable protein attribute is a distance of protein sequence from a variable reference frame.
- ⁶ 24. (Previously presented) The method of claim 20, wherein said clustering is done by Principle Component Analysis using correlation coefficient between said protein sequence-based attributes.

25. (Canceled)

- 26. (Currently amended) The method of claim 20, wherein thesaid outlier protein selected in step (e) is non-homologous to known anti-infective proteins from a pathogen selected from the group consisting of B.burgdorfei, C.jejuni, C.pneumoniae, C.trachomatis, H.influenzae, H.pylori, L.major, M.genitalium, M.pneumoniae, M.tuberculosis, N.menigitis, P.aeruginosa, P.falciparum, R.prowazekii, T.pallidum, and V.cholerae.
- 27. (Currently amended) The method of claim 20, wherein thesaid outlier protein selected in step (e) has an amino acid sequence selected from the group consisting of SEQ ID Nos: 1-31.
- 28. (Currently amended) The method of claim 20, wherein thesaid outlier protein selected in step (e) has an amino acid sequence selected from the group consisting of SEQ ID Nos: 32-118.
- 29. (Currently amended) The method of claim 20, wherein steps (a)-(c) are performed by a computer system comprising:
- (1) a central processing unit (CPU), wherein said CPU executes a program that calculates protein sequence-based attributes, wherein said protein sequence-based attributes

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comprise: percentage of charged amino acids, percentage hydrophobicity, distance of protein sequence from a fixed reference frame, measure of dipeptide complexity, and measure of hydrophobicity from a fixed reference frame; and clusters protein sequences based on said protein sequence-based attributes using Principle Component Analysis, thereby producing results;

- (2) a memory device accessed by said CPU, wherein said memory device stores said results;
 - (3) a display on which said CPU displays said results in response to user inputs; and
 - (4) a user interface device.
- 30. (Currently amended) The method of claim 20, wherein the outlier protein selected in step (e) is validated as an anti-infective by further comprising testingusing said outlier protein as for a diagnostic targetpurpose.
 - 31. (Canceled)
- 32. (Currently amended) The method of claim 20, wherein the outlier protein selected in step (e) is validated as an anti-infective by further comprising testingusing said outlier protein as a drug target for a therapeutic purpose.

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33. (Currently amended) The method-as of claim 20, wherein the outlier protein selected in step (e) is validated as an anti-infective by testing said outlier protein as a vaccine candidatewherein said outlier protein can elicit an immune response.